

26 APR 2001

FORM PTO-1449	U.S. DEPARTMENT OF COMMERCE PATENT AND TRADEMARK OFFICE	ATTY. DOCKET NO. STERN1.001APC	APPLICATION NO. Unknown 09/830703
INFORMATION DISCLOSURE STATEMENT BY APPLICANT		APPLICANT Lubbert	
(USE SEVERAL SHEETS IF NECESSARY)		FILING DATE Herewith	GROUP Unknown

U.S. PATENT DOCUMENTS

FOREIGN PATENT DOCUMENTS

EXAMINER INITIAL	DOCUMENT NUMBER	DATE	COUNTRY	CLASS	SUBCLASS	TRANSLATION	
						YES	NO
CQ	1. WO 00/31253	6/2/00	PCT				
IV	2. WO 98/59050	12/30/98	PCT				

**EXAMINER
INITIAL**

OTHER DOCUMENTS (INCLUDING AUTHOR, TITLE, DATE, PERTINENT PAGES, ETC.)

<i>CQ</i>	3.	Kessler, J., et al. Investigation of the pathogenic mechanism of parkin mutations. Society for Neuroscience Abstracts, Vol. 25, No. 1-2, 1999, p. 52-Abstract 27.20.
	4.	29th Annual Meeting of the Society for Neuroscience, Part 1, Miami Beach, Florida, October 23-28, 1999, Publication dates for the 1999 Abstract Volumes. Society for Neuroscience Abstracts, August 16, 1999.
	5.	EMBL Databases, July 13, 1999, Shimizu, N., et al., <i>Mus musculus</i> mRNA for parkin, complete cds, Abstract
	6.	Goldberg, M. S., et al., Studies of wild-type and mutant alpha-synuclein in transgenic mice, Annual Meeting Society Neuroscience, Vol. 24, No. 1/02, 1998, p. 966
	7.	Kitada, et al., Mutations in the parkin gene cause autosomal recessive juvenile parkinsonism. Nature, Vol. 392, No. 6676, April 9, 1998, p. 605-608.

EXAMINER

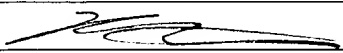
DATE CONSIDERED 1/22/02

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CQ	8. Lucking, et al., Homozygous deletions in parkin gene in European and North African families with autosomal recessive juvenile parkinsonism, The Lancet. Vol. 352, No. 9137, October 24, 1998, p. 1355-1356.
	9. Hattori, et al., Point mutations (Ihr240Arg and Ala311Stop) in the Parkin Gene, Biochemical and Biophysical Research Communications, Vol. 249, No. 3, 1998, p. 754-758.
	10. Leroy, et al., Deletions in the Parkin gene and genetic heterogeneity in a Greek family with early onset Parkinson's disease, Human Genetics, Vol. 103, No. 4, October 1998, p. 424-427.
✓	11. Abbas, et al., A wide variety of mutations in the parkin gene are responsible for autosomal recessive parkinsonism in Europe, Human Molecular Genetics, Vol. 8, No. 4, April 1999, p. 567-574.
	12. Hattori, et al., Molecular genetic analysis of a novel Parkin gene in Japanese families with autosomal recessive juvenile parkinsonism: evidence for variable homozygous deletions in the Parkin gene in affected individuals, Ann. Neurol., Vol. 44, No. 6, December 1998, p. 935-941.

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EXAMINER	DATE CONSIDERED
	1/20/02
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